(1) Publication number:

0 245 700

12

EUROPEAN PATENT APPLICATION

21 Application number: 87196197.4

(22) Date of filing: 29.04.87

(f) Int. Cl.³: **C 07 D 251/54** C 07 D 251/18, C 07 D 251/1-2 C 08 K 5/34

30 Priority: 16.05.86 US 864627

43 Date of publication of application: 19.11.87 Bulletin 87/47

Designated Contracting States: AT BE CH DE ES FR GB GR IT LI NL SE (71) Applicant: AMERICAN CYANAMID COMPANY 1937 West Main Street P.O. Box 60 Stamford Connecticut 06904-0060(US)

72 Inventor: Forgione, Peter Salvatore 120 Little Hill Road Stamford Connecticut 06905(US)

72 Inventor: Singh, Balwant 93 Janes Lane Stamford Connecticut 06905(US)

74 Representative: Wächtershäuser, Günter, Dr. Tal 29 D-8000 München 2(DE)

Alkylcarbamylmethylated aminotriazine crosslinking agents and curable compositions containing the same.

(57) Novel alkylcarbamylmethyl aminotriazines self-cure and also function as crosslinkers for compounds containing active hydrogen groups. When the active hydrogen-containing compounds are hydroxylated polymer, coatings are provided with exceptional resistance to detergent and salt-spray exposure and improved abrasion resistance. The novel aminotriazines also can be combined with fillers as binders which when cured provide shaped articles of manufacture, such as insulation and foundry core molds.

Croydon Printing Company Ltd.

ALKYLCARBAMYLMETHYLATED AMINOTRIAZINE CROSSLINKING AGENTS AND CURABLE COMPOSITIONS CONTAINING THE SAME

The present invention relates to curing agents, to curable compositions and to methods of making and using them. More particularly, the present invention relates to novel curing agents comprising alkylcarbamyl—5 methylaminotriazines and to curable compositions comprising an active hydrogen-containing material, a novel alkylcarbamylmethylaminotriazine and a cure catalyst. Coatings cured from the compositions have exceptional resistance to detergent and salt spray exposure, making them well adapted for use in powder coatings, coil coatings and can coatings. The new compositions can be used with filler to provide shaped articles of manufacture with superior properties.

BACKGROUND OF THE INVENTION

Curable compositions containing aminotriazine compounds are known in the art. As is shown in Koral et al., U.S. 3,661,819, for example, a preferred family of aminotriazine curing agents comprises (i) a triaminotriazine compound of the formula:

15

20

which will be depicted hereinafter as $C_3N_6(CH_2OR)_6$; or (ii) a benzoguanamine compound of the formula:

which will be depicted hereinafter as $C_3H_5(C_6H_5)(CH_2OR)_4$ wherein R is hydrogen or alkyl of from 1 to 12 carbon atoms. It is also known to use oligomers of such compounds, which are low molecular weight condensation products containing 5 for example two, three or four triazine rings, joined by -CH_OCH_- linkages, as well as mixtures of any of the foregoing. These are used to cure active hydrogen-containing materials, especially polymers which contain carboxyl groups, alcoholic hydroxy groups, amide groups and groups 10 convertible to such groups, such as methylol groups. such curable compositions are applied to substrates as coatings and then cured, excellent hardness, impact resistance, light stability and solvent resistance is imparted to the articles. The compositions can also be formulated with fillers and/or reinforcements such as particulate and fibrous mineral and organic materials, such as cellulose, wood, glass, graphite, textiles, silica, asbestos, wollerstonite, and the like to produce insulation, foundry molds and the like which have superior properties and show a 20 reduced tendency to emit formaldehyde during use.

As is described in German Patent OL 2,005,693

(1971) (Chemical Abstracts 76:P 34864 a (1972)), when triaminotriazines of the general formula (i) above are reacted
with arylurethanes, such as phenyl urethane, there are

25 produced reaction products of the typical formula C₃N₆(CH₂NH-COOC₆H₅)₆, and when these are reacted with polymers
containing hydroxyl groups such as acrylics and polyesters,
crosslinking occurs with the development of colorless, very
hard films, which remain colorless even when the baking time

30 is increased tenfold, to five hours at 100°C. However, subsequent experiments have shown that such coatings, like
those crosslinked with the triazines of formulae (i) and
(ii) above, are somewhat deficient in detergent resistance,
salt spray resistance and adhesion. They also are produced
35 with the liberation of phenol, which causes health and

disposal problems, and is economically wasteful.

It has now been discovered that if aminotriazines of general formulae (i) and (ii) are reacted with alkylurethanes (which are well known to be less reactive than the aryl carbamates used in the above-mentioned German Patent), derivatives are formed which are also reactive to crosslink active hydrogen-containing polymers, but the new coatings which are formed have much improved properties (detergent, salt spray, adhesion, color retention) over those of the prior art, particularly the aryl-substituted derivatives of OL 2,005,693.

Although it is known, e.g., from Amin et al., Indian J. Chem., 14B, 139-140 (1976), to prepare both aryl and alkyl carbamylmethylated melamines, by the reaction of trimethylolmelamine with n-hexyl carbamate, only the mono-substituted product was produced, and this would not be capable of acting as a crosslinker to introduce two or more urethane groups. Such groups are now believed to be essential to secure all of the advantages of the present invention.

Thus the present invention differs from the state of the art by providing aminotriazine derivatives containing at least two alkylcarbamylmethyl groups and then using them as crosslinkers for active hydrogen-containing materials to provide coatings with exceptional resistance, for example, to detergent and salt-spray exposure, and improved light stability.

SUMMARY OF THE INVENTION

According to the present invention there are provided triazine compounds selected from:

- (i) a triaminotriazine compound of the formula $C_3N_6(CH_2OR)_{6-x}(CH_2NHCOOR^1)_x$;
 - (ii) a benzoguanamine compound of the formula $C_3N_5(C_6H_5)(CH_2OR)_{4-y}(CH_2NHCOOR^1)_y$; (iii) an oligomer of (i) or of (ii); or

35

(iv) a mixture of at least two of any of (i), (ii) and (iii), wherein the R groups are, independently, hydrogen or alkyl of from 1 to 12 carbon atoms, the R1 groups are, independently, alkyl of from 1 to 20 carbon 5 atoms, x is in the range of from about 2 to about 6, and y is in the range of from about 2 to about 4.

In preferred embodiments of the invention, x is in the range of from about 2.8 to about 6 and y is in the range of from about 2.2 to about 4. With respect to compound (i) R is lower alkyl, preferably C1-C8 and R is methyl, ethyl, n-propyl, i-propyl, butyl, n-octyl, 2ethylhexyl, n-octadecyl, or a mixture of any of the foregoing. Also preferred are oligomers of (iii)(i) in which R is methyl and R^{1} is methyl, ethyl, n-propyl, i-propyl, butyl or a mixture of any of the foregoing as well as benzoguanamines (ii) wherein R and R1 are the same as defined above with respect to compound (i).

Also contemplated by the present invention are curable compositions comprising

- (a) an active hydrogen-containing material;
- (b) a triazine compound selected from
- (i) a triaminotriazine compound of the formula C₃N₆(CH₂OR)_{6-x}(CH₂NHCOOR¹)_x;
 - (ii) a benzoguanamine compound of the
- formula $C_3N_5(C_6H_5)(CH_2OR)_{4-y}(CH_2NHCOOR^1)_y$; (iii) an oligomer of (i) or of (ii) ; or

20

- (iv) a mixture of at least two of any of (i), (ii) and (iii), wherein the R groups are, indepen-
- dently, hydrogen or alkyl of from 1 to 12 carbon atoms, 30 the R1 groups are, independently, alkyl of from 1 to 20 carbon atoms, x is in the range of from about 2 to about 6, and y is in the range of from about 2 to about 4; and
 - (c) a cure catalyst.

In preferred features of this aspect of the in-35 vention, the active-hydrogen containing material (a) is a

polymeric material containing at least two reactive carboxyl, alcoholic hydroxy, amide or amine groups, or a mixture of such groups, or a group convertible to such groups, preferably a hydroxy-functional acrylic resin or a 5 low molecular weight polyester polyol. Preferably the triazine will be as set forth specifically above, and the cure catalyst will be a metal salt or metal complex comprising tin, especially preferably tetrabutyldiacetoxy stannoxane.

Alternatively, the alkylcarbamylmethyl triazines can be used as a self-crosslinkable material in providing protective and/or decorative coatings.

10

20

30

Also provided by the invention are articles of manufacture comprising substrates protectively coated with 15 a cured composition as defined above and articles of manufacture comprising a cured composition as defined above and a filler, e.g., glass, e.g., glass powder, glass beads, glass fibers or foundry sand.

DETAILED DESCRIPTION OF THE INVENTION

As starting materials to produce the alkylcarbamylmethylated triazines of this invention, there can be used the hydroxymethyl or alkoxymethyl melamines and/or benzoguanamines and oligmers thereof known in the art. Many of the starting materials are commercially available, 25 and can be made by well known procedures. In accordance with the present invention, the starting materials are reacted with alkyl carbamates, such as methyl carbamate and propyl carbamate, which also are well known in this art, in the presence of an acid catalyst.

An idealized reaction equation for the preparation of the new compounds from an alkoxymethylmelamine or a hydroxymethylmelamine is as follows:

$$c_{3}N_{6}(CH_{2}OR)_{6} + x H_{2}NCOOR^{1} \xrightarrow{H+} c_{3}N_{6}(CH_{2}OR)_{6-x}(CH_{2}NHCOOR^{1})_{x} + x ROH$$

wherein R, R¹ and x are as defined above.

The mole ratio of alkyl carbamate is selected to provide the desired degree of substitution. By way of illustration, from 2 to 6 moles can be used. Reaction is 5 typically carried out by heating in the melt or in solution, e.g., in benzene, toluene, xylene, chlorobenzene, dichlorobenzene, e.g., in the presence of catalytic amounts of acid, e.g., para-toluenesulfonic acid, nitric acid, sulfuric acid, and the like, at temperatures between 80° and 150°C., preferably 90-120°C. Measurement of the quantity of alcohol (ROH) evolved gives an indication of reaction completion. With 6 moles of alkyl carbamate, reaction is usually not 100% complete, unless forced, but a high degree of substitution, x = 5-6, is obtained. 15 Analysis by gel permeation chromotography shows that treatment of hexamethoxymethylolmelamine with substantially less than 6 moles of alkyl carbamate gives a product distribution similar to the starting material with degrees of substitution ranging up to 6. Of course, 20 only those compounds wherein at least two carbamylmethyl groups are present are crosslinkers according to this invention, even though residual alkoxymethyl groups can provide crosslinking.

Instead of alkoxymethylmelamines, hydroxymethyl25 melamines, and the corresponding benzoguanamine analogs
and oligomers can be used as starting materials. The
products can be recovered by any convenient means after
removal of byproduct water or alcohol is complete. Simply
cooling to room temperature will leave the product as a
30 residue, and the acid catalyst can be removed by neutralization.

The substituents defined by R and R¹ in the formulae above can vary widely in carbon content, and the groups can be straight chain, branched chain and alicyclic. A number of representative compounds will be exemplified in

detail hereinafter.

The active hydrogen-containing materials have as the active hydrogen group a group selected from carboxyl, alcoholic hydroxyl, amido, primary amine, secondary amine (including imine), thiol and the like. The active hydrogen-containing materials useful herein are typically film-forming compositions. Illustrative examples of active hydrogen-containing materials are shown in the above-mentioned Koral patent, the above-mentioned German OLS 2,055,693, and in Valko, U.S. 4,435,559. Typical polymers are acrylic polymers, polyesters, epoxy resins, and the like, providing that they contain active hydrogen groups.

Especially suitable are polyesters and polyacrylates containing pendant hydroxyl groups as reaction sites. 15 The former are obtained in a known manner by the reaction of polycarboxylic acids with excess quantities of polyhydric alcohols; the latter are obtained by the copolymerization of acrylic or methacrylic acid derivatives with hydroxyl-group-containing derivatives of these acids, such 20 as, for example, the hydroxyalkyl esters, optionally with the simultaneous use of additional vinyl compounds, such as, for example, styrene. Hydroxyl-group-containing polyurethanes can be obtained in known manner by the reaction of polyisocyanates with excess quantities of compounds 25 containing at least two hydroxy groups. Suitable commercially available hydroxy-group-containing polyesters are CYPLEX® 1473 and CYPLEX® 1531 from American Cyanamid Company and Cargil Polyester 5776. Suitable hydroxy- functional acrylic resins are available commercially from S.C. Johnson 30 & Son, Inc. under the trademark JONCRYL®-500. Also suitable for use are a hydroxy-terminated polycaprolactone, as well as the copolymer of 50% styrene, 20% hydroxypropyl methacrylate and 30% butyl acrylate of Example 5 of the abovementioned German OLS 2,055,693 and the polyester of phthalic 35 acid, adipic acid, ethanediol, and trimethylolpropane, with

a hydroxy number of 130 and an acid number of 1.5 of Example 6 of the said OLS publication.

As set forth herein, the curable composition includes a cure catalyst. Typically, the cure catalyst is a 5 metal salt and/or complex of a metal such as lead, zinc, iron, tin and manganese, preferably tin. Suitable salts of these metals are, for example acetates, octoates, laurates and naphthanates. Suitable complexes, for example, are tetrabutyldiacetoxy stannoxane, dibutyltin dilaurate, dimethyltin dilaurate or an acetyl acetonate. The cure catalyst is used in amounts effective to accelerate cure at the temperatures employed, e.g., 120-220°C. For example, the catalyst is used in amounts from about 0.1 to about 2.0 preferably 0.2 to 1% metal by weight (solids) based on the weight of the curable compositions.

It should also be understood that residual ether functional groups can cure with catalysts usually used with amino resins, such as acid catalysts, e.g., nitric acid, sulfuric acid, p-toluenesulfonic acid and the like.

20 This may be advantageous where lower cure temperatures are useful, e.g., when binding fillers or reinforcements, e.g., textiles, cellulose, wood flour, etc. Also useful as heterogenous acidic catalysts are ion exchange resins in the acid form.

In the practice of the invention, the curable compositions can be adapted for use in solvent-based or water based coating compositions. Coating compositions comprising aqueous dispersions are particularly suited to application by electrodeposition. Generally the compositions will contain about 1 to 75 percent by weight of resin and crosslinker combined, and the weight ratio of crosslinker to resin will range from about 5 to about 40 parts to correspondingly from 60 to 95 parts of said resin.

In many instances a pigment composition and various conventional additives such as antioxidants, surface active

agents, coupling agents, flow control additives, and the like, can be included. The pigment composition may be of any conventional type, such as, one or more pigments such as iron oxides, lead oxides, strontium chromate, carbon black, 5 titanium dioxide, talc; barium sulfate, cadmium yellow, cadmium red, chromic yellow, or the like.

After deposition on a substrate, such as a steel panel, the coating composition is devolatilized and cured at elevated temperatures by any convenient method such as 10 in baking ovens or with banks of infrared heat lamps. Curing can be obtained at temperatures in the range of from 120°C. to about 300°C., preferably from 150°C. to about 200°C. for from about 30 minutes at the lower temperatures to about 1 minute at the higher temperatures.

15

25

Conventional methods can be used to, combine the novel aminotriazines herein with fillers and/or reinforcements and to shape them into useful articles by means well known to accomplish these functions with curable aminotriazine resins. Mixing with glass fillers for example and 20 heating provides insulation shapes for pipes, and the like, after curing, and mixing with foundry sand and curing provides core molds for metal casting. These have superior strength compared to the state of the art and appear to be highly advantageous in not evolving formaldehyde during use.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The following examples illustrate the compounds and compositions of the present invention. to be construed as limiting the claims in any manner. All parts are by weight.

EXAMPLE 1

Reaction Product of 6 Moles of N-Propyl Carbamate With 1 Mole of Hexamethoxymethylmelamine (6-PC)

Hexamethoxymethylmelamine (47.2g, 0.121 mole, American Cyanamid Co. CYMEL® 300), n-propylcarbamate (75.0 g, 0.728 mole), and para-toluenesulfonic acid (0.33 g) are stirred at 95°C. in a flask equipped with a vacuum distillation head. During 50 minutes, the pressure is lowered in stages to 50 mm Hg and 21.1 g of methanol (0.659 mole, 91% of theoretical is collected in the distillate receiver. The product in the reaction flask is cooled to near room temperature, where it is a

clear, colorless, very viscous liquid. Methylene

15 chloride (100 ml) is added and stirring for one-half
hour dissolves the product. The acid catalyst is
removed by washing with sodium carbonate solution,
followed by drying over potassium carbonate. Rotary
vacuum evaporation gives 98.6 g of clear, colorless,

20 nearly solid product. Nuclear magnetic resonance (NMR) analysis shows that the product has at least five (on average) of the methoxy groups replaced by n-propyl carbamate groups:

C₃N₆(CH₂OCH₃)₀₋₁ (CH₂NHCOOCH₂CH₂CH₃)₅₋₆
25 Gel permeation chromatography shows one large peak for the monomeric compound (> 80%) and two smaller peaks corresponding to dimeric (~10%) and trimeric (~3%) oligomers.

COMPARATIVE EXAMPLE 1A

30

Reaction Product of 6 Moles of Phenyl Carbamate With 1 Mole of Hexamethoxymethylmelamine (6-PhC)

For comparison purposes, the procedure of Example 1 of German OLS 2,005,693 is repeated: Hexamethoxy-methyl-melamine (300 g, 1 mole) and 822 grams of phenyl-

urethane (6 moles) are dissolved in two liters of chlorobenzene and, after the addition of 4 grams of p-toluenesulfonic acid, are heated with a vertical condenser, with stirring and passage of CO₂, until the boiling point of chlorobenzene is reached. The methanol cleavage starts at approximately 90 to 100 C. A mixture of methanol with a small amount of chlorobenzene comes over first. With increasing temperature, the boiling point of chlorobenzene (130°C) is reached.

The mixture is then evaporated under vacuum, with stirring, to produce a colorless resin with a softening point of 85 to 120°C, which is soluble in all proportions in ethyl acetate. The yield is high. The product is of the formula $C_3N_6(CH_2NHCOOC_6H_5)_6$.

10

15

EXAMPLE 2

Reaction Product of 6 Moles of Methyl Carbamate with One Mole of Hexamethoxymethylmelamine (6-MC)

Hexamethoxymethylmelamine (19.5 g, 0.05 mole),

20 excess methyl carbamate (37.6 g, 0.50 mole), and paratoluene-sulfonic acid (0.86 g, 0.005 mole) are stirred at 97°C. in a flask equipped with a distillation head as in Example 1. The reaction mixture changes from a clear, colorless liquid to a white solid and a few ml.

25 of distillate is formed. The reaction mixture is then allowed to cool to room temperature.

A portion, 20.0 g, of the solid product is powdered and vigorously stirred with 100 ml. of water at room temperature for 1½ hours. Filtration and drying 30 gives 12.3 g of white solid; m.p. = 179-188°C. Infrared spectroscopy shows that at least 90-95% of the methoxy groups have been replaced by carbamate groups.

Purified product, 31.7 g, amounts to 98% of the theoretical yield of hexamethylcarbamylmethylated melamine, 35 a compound of the formula

C3N6(CH2NHCOOCH3)6

5

15

20

25

35

EXAMPLE 3

Reaction Product of 2 Moles of n-Octadecyl Carbamate With 1 Mole of Hexamethoxymethylmelamine

Hexamethoxymethylmelamine (25.0 g, 0.0641 mole), n-octadecyl carbamate (40.1 g, 0.128 mole), and 0.31 g

10 During 30 minutes, the pressure is lowered in stages to 50 mm Hg and distillate is collected in the receiver.

At room temperature the product is opaque, white and has the consistency of mayonnaise. Gas chromatographic analysis of a sample of the product dissolved in methyl isobutyl ketone shows practically no unconverted octadecyl carbamate. The formula is, approximately:

(0.0018 mole) of para-toluenesulfonic acid are stirred at 100°C in a flask equipped with a vacuum distillation head.

> $C_3N_6(CH_2OCH_3)_4(CH_2NCOOC_{18}H_{37}-\underline{n})_2$ EXAMPLE 4

Reaction Product of 6 Moles of Isopropyl Carbmate With 1 Mole of Hexamethoxymethylmelamine

Hexamethoxymethylmelamine (18.9 g, 0.0485 mole), isopropyl carbamate (30.0 g, 0.291 mole), and para-toluenesulfonic acid (0.13 g., 0.008 mole) and stirred at 95°C in a flask equipped with a vacuum distillation head. During 45 minutes, the pressure is lowered in stages to 50 mm Hg, and 8.46 g of methanol (0.264 mole, 91% of theoretical) is collected in the distillate receiver. is dissolved in 75 ml of methylene chloride and the solution is washed with two portions of aqueous 5% sodium carbonate to remove the acid catalyst. The solution is dried over anhydrous potassium carbonate and rotary vacuum evaporated to give 36.8 g of colorless solid (93% yield). The solid is pulverized to white powder; m.p. 101 to 130°C (clear, colorless melt). The formula is, approximately:

C₃H₆(CH₂NHCOOCH(CH₃)₂)₆ EXAMPLE 5

5

25

Reaction Product of 2 Moles of n-Octyl Carbamate With 1 Mole of Hexamethoxymethylmelamine

Hexamethoxymethylmelamine (22.6 g, 0.0579 mole), n-octyl carbamate (20.0 g, 0.116 mole), and para-toluenesulfonic acid (0.19 g, 0.0011 mole) are stirred at 75°C in a flask equipped with a vacuum distillation head. 10 50 minutes, the pressure is lowered in stages to 50 mm Hg, and 3.58 g of methanol (0.112 mole, 96% of theoretical) is collected in the distillate receiver. The product is dissolved in 150 ml of methylene chloride and the solution is washed with two portions of aqueous 5% sodium carbonate 15 to remove the acid catalyst. The solution is dried over anhydrous potassium carbonate and rotary vacuum evaporated to give 37.2 g of almost-clear, colorless, viscous liquid (96% yield of theoretical product). The infrared spectrum is consistent with the expected structure. Gel permeation 20 chromatography shows peaks attributed to mono-, di-, tri-, and higher-substituted products; little oligomeric material is evident. The formula is, approximately:

$C_3N_6(CH_2OCH_3)_4(CH_2NHCOOC_8H_{17}-\underline{n})_2$ EXAMPLE 6

Reaction Product of 6 Moles of n-Propyl Carbamate With 1 Mole of Hexamethylolmelamine

The general procedure of Example 1 is repeated, substituting the hydroxymethyltriazine: Hexamethylol
melamine (10.0 g, 0.0327 mole), n-propyl carbamate (20.2 g, 0.196 mole), and para-toluenesulfonic acid (0.60 g, 0.0035 mole) are stirred at 95°C in a flask equipped with a vacuum distillation head. During 30 minutes, the pressure is lowered in stages to 50 mm Hg., and 3.24 g. of distillate (mostly water, 92% of theoretical) is collected

in the distillate receiver. At room temperature, the product in the reaction vessel is a gray-white solid. The infrared spectrum shows little or no residual hydroxyl functionality and is similar to the spectrum of authentic bexa-n-propylcarbamylmethylated melamine, made by Example 1. The general formula is, approximately:

C₃N₆(CH₂NHCOOC₃H₇-<u>n</u>)₆

10

20

25

30

EXAMPLE 7

Reaction Product of 3.5 Moles of n-Propyl Carbamate with 1 Mole of Hexamethoxymethylmelamine

The general procedure of the preceding Examples is used to react hexamethoxymethylmelamine (167.8 g, 0.430 moles) with 155.1 g, 1.566 moles of n-propyl carbamate. The acid catalyst in this instance is 0.7 g of concentrated nitric acid. The product weighs 254.2 g and contains only 0.1% residual carbamate. It melts at 85-95°C. A 65% solids solution in xylene remains clear and colorless for 8 weeks. It has the formula, approximately:

 $C_{3}^{N_{6}}(CH_{2}^{OCH_{3}})_{2.5}(CH_{2}^{NHCOOC_{3}H_{7}-\underline{n}})_{3.5}$

Reaction Product of 3.5 Moles of Methyl Carbamate With 1 Mole of Hexamethoxymethylmelamine

The procedure of Example 7 is repeated substituting 1.589 moles of methyl carbamate. The product weighs 241.7 g and melts at 95-103°C. It has the formula, approximately:

C₃N₆(CH₂OCH₃)_{2.5}(CH₂NHCOOCH₃)_{3.5} EXAMPLE 9

> Reaction Product of 3.5 Moles of n-Propyl Carbamate With I Mole of Hexamethoxymethylmelamine Oligomer

The procedure of Example 7 is repeated, substi-35 tuting an oligomeric triazine (American Cyanamid Co. CYMEL® 303), $C_3N_6(CH_2OCH_3)_{5.25}$. The catalyst is removed by extracting a xylene solution of the product mixture with sodium carbonate solution. A 75% solution in xylenes remains clear and colorless for more than 6 weeks. The product melts at 74-80°C. The formula is, approximately:

C₃N₆(CH₂OCH₃)_{1.75}(CH₂NHCO₂C₃H₇)_{3.5} EXAMPLE 10

10

20

30

Reaction Product of 4 Moles of n-Propyl Carbamate With 1 Mole of Hexamethoxymethylmelamine Oligomer

The procedure of Example 9 is repeated with the higher mole ratio of n- propyl carbamate. The product, 412.2 g, melts at 80-90°C. At 75% solids in xylenes and methyl isobutyl ketone, the product in solution remains clear and colorless for more than 6 weeks. It has the following approximate formula:

C₃N₆(CH₂OCH₃)_{1.25}(CH₂NHCOOC₃H₇-n)_{4.0} EXAMPLE 11

EXAMPLE II

Reaction Product of 5 Moles of n-Propyl Carbamate With 1 Mole of Hexamethoxymethylmelamine Oligomer

The procedure of Example 9 is repeated with a higher mole ratio of \underline{n} -propyl carbamate. The product, 32.4 g, melts at 80-92°C. It has the following approximate formula:

C₃N₆(CH₂OCH₃)_{0.25}(CH₂NHCOOC₃H₇-n)_{5.0} EXAMPLE 12

Reaction Product of 4 Moles of n-Propyl Carbamate and 2 Moles of Methyl Carbamate With 1 Mole of Hexamethoxymethylmelamine

The general procedure of Example 9 is used at 95°C. The product, 95.8 g, melts at 85-95°C. The formula is, approximately:

 $C_3^{N_6}(CH_2^{NH-CO_2}CH_3)_{1.25}(CH_2^{NHCO_2}C_3^{H_7-n})_4$

EXAMPLE 13

Reaction Product of 5.25 Moles of n-Propyl Carbamate With 1 Mole of Hexamethoxymethylmelamine Oligomer

5 The general procedure of Example 9 is used. The product weighs 113.5 g and melts at 90-100°C. The formula is, approximately:

C₃N₆(CH₂OCH₃)_{0.15}(CH₂NHCOOC₃H₇-n)_{5.1} <u>EXAMPLE 14</u>

Reaction Product of 2.5 Mole of n-Propyl Carbamate and 2.5 Mole of Methyl Carbamate With 1.0 Mole of Hexamethoxymethylmelamine Oligomer

10

20

The general procedure of Example 9 is used at 110°C. The product weighs 174.5 g and melts at 85-95°C. The general formula is, approximately:

C₃N₆(CH₂OCH₃)_{0.25}(CH₂NHCO₂CH₃)_{2.5}(CH₂NHCO₂C₃H₇-n)_{2.5} EXAMPLE 15

Reaction Product of 3.5 Mole of n-Propyl Carbamate with 1 Mole of (Trimethoxy-methyl-Tributoxymethyl) Melamine

(Trimethoxymethyl-tributoxymethyl) melamine (CYMEL® 1133, American Cyanamid Co., 320.3 g, 0.689 mole), n-propyl carbamate (248.3 g, 2.411 mole) and 1.12 g of concentrated nitric acid are stirred and heated in a 100°C. oil bath under a steady stream of nitrogen. Distillate is collected in a dry-ice/isopropanol cooled After 60 minutes, during which vacuum of up to 50 mm Hg is applied, the reaction is stopped. The hot, crude reaction product is dissolved to 30% solids in mixed xylenes (1059 g of xylenes are added) and extracted once with 300 ml of 5% aqueous sodium carbonate solution. The organic layer is then extracted several times with hot deionized water to neutralize the acid and reduce the amount of residual n-propyl carbamate. 35 The organic layer is dried over potassium carbonate

(anhydrous) until clear and then stripped under vacuum of a 65% solids content. The product weighs 698.5 grams.

5

15

EXAMPLE 16

Reaction Product of 3.5 Mole of Methyl Carbamate with 1 Mole of (Trimethoxymethyl-Tributoxymethyl)Melamine

The procedure of Example 15 is repeated, substituting 376.28 g, 0.809 mole of trimethoxymethyl
tributoxymethylmelamine, 212.42 g, 2.832 mole of methyl carbamate and 1.32 g of conc. nitric acid. The product is recovered by the procedure of Example 9.

In the following examples, the alkylcarbamylmethylated triazines of this invention are formulated
into curable compositions and evaluated as coatings.
For comparison purposes, the phenylcarbamylmethylated
triazine of German OLS 2,005,693 (Example 1A herein) is
also evaluated.

The general method of preparation is as follows: 20 Thermosetting coatings containing polyols with alkyl and phenylcarbamylmethylated melamines on steel are prepared by mixing high solids solutions of alkylcarbamyl and arylcarbamylmethyl melamines (65% solids in methyl isobutyl ketone or xylene) with a solution of a hydroxyl 25 functional acrylic resin (specifically, JONCRYL®-500, S.C. Johnson & Son, Inc.; 85% solids in methyl amyl ketone), a solution of a tin catalyst (tetrabutyldiacetoxy stannoxane, TBDSA, 10% solids in methyl isobutyl ketone), and a flow control additive (FC-431, 10% solids in ethyl acetate, 3M 30 Co.). A second series of coatings compositions is prepared as above but with low molecular polyester polyol as backbone resin (specifically, CYPLEX® 1473, 60% solids in Xylene, American Cyanamid Co.). Both systems are formulated with and without EPON®-1001, an epoxy resin (10

parts per hundred-phr, as an 85% solids solution in methyl isobutyl ketone - Shell Co.) to assess resistance properties in particular.

The coatings are applied using #40 or #46 WIRE
5 CATORS® by drawdown of resin formulations on 4" x 12"

BONDERITE®-100 treated steel panels. The panels, after drawdown, are held for 10 minutes at room temperature and then cured on racks in a temperature controlled oven, at specified temperatures. The coatings prepared are about 1.2 ± 0.2 mils thick and initially tested for solvent resistance by rubbing with a cloth saturated with methyl ethyl ketone (MEK rubs) in accordance with standard testing methods.

Table 1 illustrates a typical charge composition 15 for preparing a coated panel.

	TABLE 1									
	<u>Material</u>	% Solids	PHR*	Charge (Grams	solution)					
	CYPLEX® 1475-5	65	70	32.3						
	Propyl Carbamate	•	•							
20	Melamine resin	60	20	10.0						
	EPON-1001	75	10	4.0						
	TBDAS cat.	10	1	3.0						
	FC-431	10	0.13	0.4						

^{*} PHR = parts per hundred resin; final solution 57.8% solids.

25 Materials 1--5 are stirred until homogeneous, then filtered through a 10 micron felt filter to remove small particles and deaerated.

The properties of the coatings evaluated include:

Method Property Forward and Reverse Impact ASTM D-3281-73 Visual Measured on Glossgard II 5 20° Gloss 20°/60° Gossmeter - Neotec Instr. Div., Pacific Scientific Detergent Resistance at 72°C. ASTM D-2248-73; reapproved 10 1982; Evaluation of Degree of Blistering of Paints D-714 Blister Classification Example: F-8 means few small blisters; D-4 means dense large blisters. The smaller the number following the 15 letter, the larger the blister on a scale of 1-10, with 10 meaning no blistering.

EXAMPLES 17-22

The crosslinker of Example 1 herein, the reaction product of 6 moles of n-propyl carbamate and 1 mole of hexamethoxymethyl melamine (6-PC), is used with a hydroxy-functional polyacrylate and tetrabutyldiacetoxystannoxane as cure catalyst. For comparison purposes, fomulations are made substituting the reaction product of 6 moles of phenyl carbamate with hexamethoxymethyl melamine (6-PhC) of Comparative Example 1A. The formulations used and the properties of the cured films are set forth in Table 2:

TABLE 2: Carbamylmethylated Melamine Crosslinked Acrylic Resin

	Crosslinke	d Acryl	Lic Resi	n			
	Example	17	18	19	20	21	22
	Composition (parts)	y weigh	it)				
5	Polyacrylatea	80	70	60	50	80	60
	6-PC ^b	20	20	40	40	20	40
	6-PhC ^C						
	Epoxy Resin ^d		10		10		
	TBDAS ^e .	1	1	1	1	1	1
10	Cure Temp., °C.	180	180	180	180	160	160
	Cure Time, min.	20	20	20	20	30	30
	Properties				•		
	MEK wipes	200+	200+	200+	200+	200+	200+
	Thickness, mils.	1.2	1.1	1.0	1.0	1.2	1.1
15	Forward Impact,	30	10	10	20	20	
	Color	Clr	Clr	Clr	Clr	Clr	Clr
	20° Gloss	97	93	95	93	97	96
	Knoop Hardness	13.5	13	16	15.8	10.8	16.2
	Detergent Immersion						
20	72 hrs	8M	10	F9	10	8M	D9
	120 hrs	D8	10	F8	10	Мб	D8
	192 hrs	D8	D9	F9	10	D7 .	м9
	240 hrs	-			10		
	288 hrs '				10		
25	408 hrs				10		
	456 hrs				10		
	744 hrs				10		
	912 hrs				10		
	1100 hrs				10		

	TABLE 2: CONTINUED						
	Example	17	<u>18</u>	19	20	21	22
	Composition (parts b	y wei	ght)				
	Polyacrylatea	80	70	60	50	80	60
5	6-PC ^b		~~	***			
	6-PhC ^C	20	20	40	40	20	40
	Epoxy Resin ^d		10		10		
	TBDAS ^e	1	1	1	I	1	ı
	Cure Temp., °C.	180	180	180	180	160	160
10	Cure Time, min.	20	20	20	20	30	30
	Properties						
	MEK wipes	Tend 4mb	200 ^m	200+	200+	200 ^m	~
	Thickness, mils.	1.1	0.94	0.88	0.83	1.2	1.0
	Forward Impact,	~	20	· 20	20	10	
15	Color	Clr	Yel	Sl.Brn	Yel	Sl.Yel	Clr
	20° Gloss	94	97	95	91	92	86
•	Knoop Hardness	15.3	13	18	17	12.4	19.4
	Detergent Immersion		•				
	72 hrs	Мб	10	F6	10	VD6	M8.
20	120 hrs	D2	М8	F8	10	VD4	M8
	192 hrs		D8	F8	10	~-	M8
	240 hrs				10		
	288 hrs ·		1		10		
	408 hrs				10		
25	456 hrs				F6	-:-	
	744 hrs				F6		
	912 hrs .				F6		
	1100 hrs				F6	~~	

JONCRYL® S.C. Johnson & Son, Inc., 85% solids in methylamylketone;
b n-propylcarbamylmethyl melamine (Example 1);
c phenylcarbamylmethyl melamine (Comp. Ex. 1A);
d EPONO 1001, Shell Polymers Inc.;
e Tetrabutyldiacetoxy stannoxane;
35 m = marred; and s = softened

EXAMPLES 23-30

The crosslinkers of Examples 7 and 8 herein (3.5 PC and 3.5 MC) are used to crosslink the hydroxy functional acrylic resin and evaluated in coatings. The formulations used and the properties obtained are set forth in Table 3:

•	TABLE 3: Carbamy	lmethyl	ated M	<u> </u>	ne-Cros	slinke	ed Acry	/lic Re	sins
	Example	23	24	25	26	27	28	. 29	30
	Composition (par	ts by w	reight)	1					
	Polyacrylatea	80	70	60	50	80	70	60	50
5	3.5-PC ^b	20	20	40	40				
	3.5-MC ^C					20	20	40	40
	Epoxy Resin ^d		10		10		10		10
	TBDAS e	1	1	1	1	·1	1	1	1
	Cure Temp., °C.	180	180	180	180	160	160	180	180
10	Cure Time, min.	20	20	20	20	20	20	20	20
	Properties								
	MEK wipes	200+	200+	200+	200+	200+	200 ^m	200+	200
	Thickness, mils.	1.0	1.1	1.2	1.1	1.0	1.0	0.95	0.96
	Forward Impact,	30	5	20	18.5	30	15	15	20
15	Color	Clr	Clr	Clr	Clr	Clr	Clr	Clr	Clr
	20° Gloss	90	95	95	91	97.5	99.5	101	101
	Knoop Hardness	11.8	10.4	15.4	18.5	13	15	16.4	17
	Detergent Immer	sion'							
	48 hrs	8M	65 es						
20	72 hrs		10	M8	10	М9	10	10	10
	120 hrs	D8	D8	M4	10	D6	10	10	10
	192 hrs		D8	D8	10	D8	10	10	10
	240 hrs		***		10		м8	F8	10
	288 hrs	~~	-		10	~-		F8	10
25	408 hrs				10	~		Мб	10
	456 hrs			-	10			M4	10
	744 hrs	~-			10			M4	10
	912 hrs				10	~~ ~	****	M4	10
	1100 hrs	~-			10		***		10
30	1400 hrs				10	***			F9

þ

See footnote Table 2
3.5 propyl carbamylmethylmelamine (Ex. 7)
3.5 methyl carbamyl methylmelamine (Ex. 8)
See footnote Table 2
-do-

c d

³⁵ e

EXAMPLES 31-34

The crosslinker of Example 1 herein, the reaction product of 6 moles of n-propyl carbamate and 1 mole of hexamethoxymethylolmelamine (6-PC) is used with a low 5 molecular weight hydroxyfunctional polyester and tetrabutyldiacetoxy stannoxane as cure catalyst. For comparison purposes, formulations are made substituting the reaction product of 6 moles of phenylcarbamate with hexamethoxymethylmelamine (6-PhC) of Comparative Example 1A. The formulations used and the properties of the cured films are set forth in Table 4:

TABLE 4: Carbamylmethylated Melamine-Crosslinked Polyester Resin

	Example	31	32	33	34	31A
	Composition (parts by					
5	Polyestera	80	70	60	50	80
	6-PC ^b	20	20	40	40	
	6-PhC ^C					20
	Epoxy Resin ^d		10		10	
	TBDAS ^e	1	ı	1	1	1
10	Cure Temp., °C.	180	180	180	180	180
	Cure Time, min.	20	20	20	20	20
	Properties					•
	MEK wipes	200 ⁸	200+	200	200+	200 ⁸
	Thickness, mils.	.98	1.0	.90	1.0	.92
15	Reverse impact,	160		40		160
	Forward Impact,	160		70		160
	Color	Clr	Clr	Clr	Clr	Clr/Yel
	20° Gloss	99	102	104	102	92
	Knoop Hardness	15.4	18.8	19.5	18.8	14.6
20	Detergent Immersion	,			•	
	48 hrs		10		10	
	72 hrs	10		10		10
	96 hrs	10		10		M8
	168 hrs		10	10	10	
25	216 hrs	Мб	10	F8	10	D4
	336 hrs		M8		10	
	398 hrs	M2		M4		
	504 hrs		D4-6	M2	10	
	652 hrs				F8	

		TABLE 4:	CONTIN	UED		
	Example	321	<u>33A</u>	34A	34E	34C
	Composition (parts	by weight	<u>t)</u>			
	Polyestera	_ 70	60	50	80	. 60
5	6-PC ^b					
•	6-PhC ^C	. 20	40	40	40	40
	Epoxy Resin ^d	10		10		
	TBDAS ^e	1	1	1	1	1
	Cure Temp., °C.	180	180	180	160	160
10	Cure Time, min.	[*] 20	20	20	30	30
	Properties					
	MEK wipes		200	200	200	200 ^m
	Thickness, mils.	1.1	. 85	.85	.95	.65
	Reverse impact,		140	30	160	160
15	Forward Impact,		150	70	160	160
	Color	Clr	Yel	SlYel	Clr	Clr
	20° Gloss	106	107	105	98	101
	Knoop Hardness	19.1	21	22.5	16.2	· 21
	Detergent Immersio	n				
20	48 hrs					en -
	72 hrs	10	8M	10	10	F8
	96 hrs		D8	10	10	М6
	168 hrs			10		
	216 hrs	. 10	D4	F8	F9	D8
25	336 hrs			****		
	398 hrs	10		D4	D2	D2
	504 hrs					
	652 hrs	. D8				

a American Cyanamid Co., CYPLEX® 1473-5

³⁰ b-f See footnote above, Table 2.

EXAMPLES 35-44

The crosslinkers of Examples 7 and 8, (3.5-PC and 3.5-MC) are used with the low molecular weight hydroxyfunctional polyester and tetrabutyldiacetoxy5 stannoxane as cure catalyst. The formulations used and the results obtained are set forth in Table 5:

Ì

TABLE 5: Carbamylmethylated Melamine-Crosslinked Polyester Resins

	POLYES	rer vear	15			
	Example	35	36	37	38	<u>39</u>
5	Composition (parts by	weight)	_			
	Polyester ^a	80	70	60	50	80
	3.5-PC ^b	20	20	40	40	
	3.5-MC ^C					20
	Epoxy Resin ^d		10		10	
10	tedas ^e	1	1	1	1	1
	Cure Temp., °C.	180	180	180	180	180
	Cure Time, min.	20	20	20	20	20
	Properties			•		
	MEK wipes	200 ^m	200±	200 ^S	200÷	200÷
15	Thickness, mils.	.92	•9*	.77 ^	.96	-88
	Reverse impact,	160		120		150
	Forward Impact,	160	100	120	70	160
	Color	Clr	Clr	Clr	Sl.Yl.	Clr
	20° Gloss	96	85	95	90	99
20	Knoop Hardness	14.2	17.4	18.8	21	14.4
	Detergent Immersion					
	48 hrs	10	10			
	72 hrs			10	10	10
	96 hrs			10	10	M8
25	168 hrs	M6	10			
	216 hrs	D2	M8	мб	10	D4
	336 hrs		D2		****	'
	398 hrs			D2	F8	
	504 hrs				F6	

		TABLE 5:	CONTINU	JED		
	Example	40	41	42	43	44
	Composition (parts	by weight)				
	Polyestera	70	60	50	80	60
5	3.5-PC ^b					
	3.5-MC ^C	20	40	40	20	40
	Epoxy Resin ^d	10	-	10	-~	
	TBDAS ^e	1	1	1	1 .	1
	Cure Temp., °C.	180	180	180	160	160
10	Cure Time, min.	20	20	20	30	30
	Properties					•
	MEK wipes	200+	200+	200+		
	Thickness, mils.	1.2	.85	1.1	1.1	1.0
	Reverse impact,	50	90	10		
15	Forward Impact,	50	80	40		
	Color	sl.Yl.	Clr	Sl.Yl.	Clr	Clr
	20° Gloss	101	99	102	101	102
	Knoop Hardness	20	22.5	23.5	18	17.2
	Detergent Immersion					. 50
20	48 hrs					
	72 hrs	10	10	10	10	10
	96 hrs	10	M8	10	10	10
	168 hrs					
	216 hrs	10	D4	10	F8	10
25	336 hrs					10
	398 hrs	10		10	M4	haze
	504 hrs			10		VD8
	652 hrs			VF8	VD2	s

See footnote above, Table 4
-do- , Table 3
-do- , Table 3 30 þ

c d

⁻do-

e -do-

m

⁻do-35

A review of the data in the foregoing tables indicates that improved properties are obtained with the alkylcarbamylmethylated melamines of this invention in comparison with the phenylcarbamylmethylated melamine of 5 the prior art. Particularly outstanding with the acrylic and polyester coatings are the excellent detergent resistance properties of the new alkylcarbamylmethylated melamines. For example, both the methyl and propyl carbamylmethylated melamines give over 1100 hours of blister free 10 coatings in a detergent bath when formulated with acrylic polyol and Epon-1001 versus 408 hours for the phenyl system. In the polyester system, the alkylcarbamate system gives 504 hours of blister free coatings versus 398 hours for the phenylcarbamate system in a detergent bath. 15

EXAMPLES 45-52

Unpigmented coating formulations are prepared by the general procedure described above and cured on steel panels at 177°C. for 6-PC and benzoguanamine resins

(CYMEL® 1123) and at 125°C. for melamine oligomer methoxymethyl (CYMEL® 303) resins, using hydroxy functional polyesters and 20 min. cure times. In addition to detergent resistance, salt spray resistance and pencil hardness are measured. The formulations used and the results obtained are set forth in Table 6, as follows:

	Table 6. Carbamylme	thylme	lamine-	Cured P	olyest	ers
	Example	45	46	47	48	49
	Composition (parts by wei	ght)				
	Polyester (CYPLEX®1473-5)	60	- 70	75	80	85
5	6-PC	40	30	25	-20	15
	CYMEL® 303		•			
	CYMEL® 1123					
	EPON® 1001					
	TBDAS	1	1	1	1	1
10	p-TSA					
	Properties					
	Pencil hardness	3H-4H	2H-3H	2H-3H	3H	2H-3H
	Knoop hardness	23	16.4	16.6	17.5	16
	Reverse Impact	60	150	150	160	160
15	T-Bond	T 5	T2	T4	тз	Tl
	Detergent Resistance,					
	Blister Code, Hrs.					
	48	F9	F 8	м9	м9	8M
	72	МŚ	M8	M8	м8	м8
20	216	D7.	D6	. D7	D7	D6
	384	D 7	D4	D6	D6	D4
	Salt Spray Exposure					
	Blister Code, 46 Hrs.	10	10	10	10	F8
	Tape Pull, mm	0	· 1	1	1	0
25	168 hrs	10	10	10	10	10
	mm	1	2	2	2	3
	300 hrs	10	10	10	10	10
	mm	2.5	. 2	2.5	2	. 4
	468 hrs	10	10	10	10	10
30	mm	4	3.5	4	3.5	5

		-32-				
	•			•		
	Table	e 6. (C	ontinue	<u>a)</u>		
	Example	50	<u>51</u>	52	50A	51A
	Composition (parts by we	ight)				
	Polyester (CYPLEX®1473-5)) - 75	70	65	75	75
5	6-PC	15	20	25		
	CYMEL® 303				15	
	CYMEL® 1123					15
	EPON® 1001	10	10	10	10	10
	TBDAS	1	1	1		
10	p-TSA ·				.4	.4
	Properties					
	Pencil hardness	2H-3H	3H-4H	4H-5H	2H-3H	H-2H
	Knoop hardness	₌ 16	21.5	19.4	15.6	15.5
	Reverse Impact	160	160	160	160	150
15	T-Bond	T2	T4	T4	T2	T4
	Detergent Resistance,					
	Blister Code, Hrs.					
	48	10	10	10	10	10
	72	10	10	10	<u>,</u> 10	10
20	216	10	F9	10	(168-D9)	D7
	. 384	10	D9	10		D4
	Salt Spray Exposure					
	Blister Code, 46 Hrs.	10	10	10	10 .	10
	Tape Pull, mm	. 0	0	0	o .	0
25	168 hrs	10	10	10		10
	mm	0	Ο.	· 0		0
	300 hrs	10	10	10		10
	mm	- 0	Ö	0		0
	468 hrs	10	10	10		
30	- mm	0	0	0		
	1080 hrs	10	10	10	D4	F2
	mm .	0	0	0	0	4
	1440 hrs	10	10	10	M4/6	·
	mm	. 0	0	0	Striped	

EXAMPLES 53-60

Unpigmented coating formulations are prepared by the general procedure described above and cured on steel panels at 177°C. for 6-PC and at 125°C. for the melamine oligomer resin, using a different hydroxyfunctional polyester and a 20 min. cure time. The formulations used and the results obtained are set forth in Table 7, as follows:

	Table 7. Carbamylm	ethylm	elamine-	Cured E	olyeste	rs
	Example	53	. <u>54</u> .	<u>55</u>	56	57
•	Composition (parts by we	ight)	.			
	Polyester (CYPLEX@1473-5) · 60	67	70	75	80
5	6-PC	40	33	30	25	20
	CYMEL® 303					
	EPON® 1001			~-		
	TBDAS	1	1	1	1	1
	p-TSA	·				
10	Properties					
	Pencil hardness	4H	4H	4H	2H-3H	2H-3H
	Knoop hardness	16.6	16.4	16.4	15.2	14.2
	Reverse Impact	40	110	120	160	160
	T-Bond	Т4	т3	Т3	Т3	T2
15	Detergent Resistance,		٠.			
	Blister Code, Hrs.					
	48	F9	F9	F9	м9	F9
	72	F9	F8	F9	M8	F9
	216	D6	D6	D7	D8	BQ
20	384	M/D7	D5	D7	D7	D7
	Salt Spray Exposure					
	Blister Code, 46 Hrs.	10	10	10	10	10
	Tape Pull, mm	0	1	0	2	1
	168 hrs	10	10	10	10	10
25	mm	0	2	2	3	2
	300 hrs	10	10	10	10	10
	min	0	2	4	4	4
	468 hrs	10	10	10	10	10
	mm	1	4	5	6	6
30	1080 hrs					
	mm					
	1440 hrs					
	nun					

:,

•	Table 7. (Continued)									
	Example	<u>58</u>	59	60	58A	58B				
	Composition (parts by wei	ght)								
	Polyester (CYPLEX®1473-5)	75	70	65	75	75				
5	6-PC	15	20	25						
	CYMEL® 303				25	15				
	EPON® 1001	10	10	10	***	10				
	TBDAS	1	1	1						
	p-TSA				0.4	0.4				
10	Properties									
	Pencil hardness	3н-4н	4H	3H-4H	H-2H	2н-3н				
	Knoop hardness	15.4	16.6	15.5	12.6	14				
	Reverse Impact z	160	160	160	150	130				
	T-Bond	T2	Т3	T4	TЗ	T 5				
15	Detergent Resistance,									
	Blister Code, Hrs.									
	. 48	10	10	10	10	F8				
	72	10	10	10	F9	F8				
	216	8M	M9	F9	D7	D6				
20	384	M8	M/D9	D9	D4					
	Salt Spray Exposure									
	Blister Code, 46 Hrs.	10	10	10	10	D8				
	Tape Pull, mm	0	0	0	0	4				
	168 hrs	10	10	10	D9	~-				
25	mm.	0	0	0	2	5				
	300 hrs	10	10	10		~-				
	mm	0	0	0	3	8				
	468 hrs	10	10	10		~-				
	mm	0	0	0	4	10				
30	1080 hrs	10	10	. 10	Ml					
	nun	0	0	0	disint.	disint.				
	1440 hrs	10	M /	10		· ·				
	mm	1	0	0						

-36-

EXAMPLES 61-68

Titanium dioxide pigmented coating formulation are prepared by the general procedure described above and cured on steel panels, using 6-PC and methylolmel-.

5 amine and methylolbenzoguanamine resins, the latter two as controls. The formulations used and the results obtained are set forth in Table 8 as follows:

Table 8. Tio	22-	- Pigm	ented	Cured	Coating	S
	_		~ ~			_

	14D16 0. 110	22	enceu	Carea	CORCE	93	
	Example	61A	61	62	62A	63A	63
	Composition (parts by	weight)					•
	JONCRYL®500ª		~	75	75		
5	AROPLAZ 1710 R60 ^b	75	75				
	CYMEL® 1473-5°		~				
	CARGILL 5775 ^d					75	75
	CYMEL® 303 ^C	25				15	
	CYMEL® 1123 ^C				25		
10	CYMEL 6-PC ^e		25	25			25
	EPON® 1001 ^f					10	
	T-12 ^g						
	TBDAS ^h		1.	ì			1
	p-TSA ⁱ	0.4			0.5	0.4	
15	Cure Temp °C.	125	177	177	177	125	177
	Properties						
	MEK Resist	200+	200+	200+	200+	200	200+
	Cross-Hatch Adhesion	3	5	4	5	5	5
	Knoop hardness	16.6	17.5	26	18	10.8	21
20	Reverse Impact	5	30-40	5	5	160	40
	Film Thickness, mils	1.2	1.1	1.1	1.3	1.3	1.2

a S. C. Johnson & Son, Inc., hydroxyfunctional polyacrylate;

b Spencer Kellog, Div. of Textron, Inc., siliconized polyester;

²⁵ c American Cyanamid Company;

d Cargill Co.

e Hexa(propylcarbamylmethyl)melamine (Ex. 1);

f Shell Chemcial Co.;

g Dibutyltin dilaurate;

^{30.} h Tetrabutyldiacetoxy stannoxane;

i p-toluenesulfonic acid;

Table 8. (Continued)								
	Example	61A	61	<u>62</u>	62A	63A	63	
	Detergent Resistance,							
	Blister Code, Hrs.					•		
5	48	10	10	10	D8	10	Ф9	
•	96	M/D8/9	D8	10	D6	м9	р9	
	124			10	D9	M		
	148			10	YD6	VD9	VD9	
	192	D8	D6	10	D4	peeled	VD9	
10	288	VD6	D4	D9	D2		VD4	
	336			D9	D2		•	
	Salt Spray Exposure							
	Blister Code, 46 Hrs	. 10	10	10	D9	10 ,	10	
	Tape Pull, mm	1	0	0	O _.	0	0.5	
15	336 hrs	10	D8	10	D9	10	10	
	mm	2	1.5	0	3	1	0	
	800 hrs	10	D8	10	D9	10	10	
	mm	5	5	0	11	8	0	

NC = no change

The foregoing examples show that even without the epoxy resin (EPON® 1001) additive, the alkylcarbamate melamine systems outperform all state of the art melamine systems, which include the phenyl carbamate, alkoxymethyl and benzoguanamine systems in detergent resistance. The above-mentioned results are unexpected because alkyl urethanes on heating are known to be poorer leaving groups than the phenoxy blocked isocyanate system and therefore they would be expected to give somewhat inferior coating properties. The foregoing examples also show that the alkylcarbamylmethylated melamines of this invention also provide coatings with outstanding salt spray resistance in comparison with other melamine based systems.

It is generally the case also, that the alkylcarbamylmethylated melamines of this invention afford coatings
with good color stability. While the phenyl analog gives
off white to tan coatings on cure, the alkyl systems are
unchanged. In addition, the phenylcarbamate based coatings
on exposure to U.V. light change to a much darker color
(tan to light brown), while the alkylcarbamate systems
change only slightly to a light tan color. Finally, the
data show that while outstanding resistance properties and
color stability have been obtained with the alkylcarbamate
melamines, other important and desirable coatings properties such as Knoop hardness, impact and solvent resistance
(MEK rubs + 200) have been maintained as is the case with
conventional resins.

EXAMPLE 69

The product of Example 1 can be used as a binder 30 for foundry sand. The binder as an acetone solution (e.g., 77 wt. %) containing 1 part/100 of TBDSA catalyst is kneaded with sand and the solvent is vaporized by heating. The amount of binder to sand is preferably 1-5.5 parts to 100 parts of sand. The coated sand is then filled into a core mold and heated at 200-300°C. for 30 sec. to two

minutes. A core having good strength and showing little tendency to give off gases having a strong smell (e.g., formaldehyde) will be obtained.

Instead of sand, glass powder and glass fiber can 5 be substituted, in which case, thermally insulating shapes having good structural integrity will be obtained.

The above-mentioned patents and publications are incorporated herein by reference. Many variations of this invention will suggest themselves to those skilled in this 10 art in light of the above, detailed description. of using n-propylcarbamylmethylated- and methylcarbamylmethylated melamines as curing agents in the formulations of Tables 1-8, the corresponding alkyl (and mixed alkyl) carbamylmethylated melamine and melamine oligomers of 15 Examples 3-16 can be used. Instead of tetrabutyldiacetoxy stannoxane and, dibutyltin dilaurate as cure catalysts, lead octoate, and stannous octoate can be used. Instead of hydroxyfunctional polyesters and polyacrylates, epoxy resins, such as the polyglycidylethers of bisphenol A and 20 the reaction products thereof with amines and ammonia can be used. All such obvious modifications are within the full intended scope of the appended claims.

122 CLAIMS:

- 1. A triazine compound selected from
- (i) a triaminotriazine compound of the formula $C_3N_6(CH_2OR)_{6-x}(CH_2NHCOOR^1)_x$;
- (ii) a benzoguanamine compound of the formula $C_3N_5(C_6H_5)(CH_2OR)_{4-y}(CH_2NHCOOR^1)_y$;
- (iii) an oligomer of (i) or of (ii); and
- (iv) a mixture of at least two of any of (i), (ii) and (iii), wherein the R groups are, independently, hydrogen or alkyl from 1 to 12 carbon atoms, the R¹ groups are, independently, alkyl of from 1 to 20 carbon atoms, x is in the range of from about 2 to about 6, and y is in the range of from about 2 to about 4.
- 2. A triaminotriazine compound (i) as defined in Claim 1 wherein x is from about 2.8 to about 6, and wherein R is C_1 - C_8 alkyl and R^1 is methyl, ethyl, <u>n</u>-propyl, butyl, <u>i</u>-propyl, <u>n</u>-octyl, 2-ethylhexyl, <u>n</u>-octadecyl, or a mixture of any of the foregoing.
- 3. A benzoguanamine compound (ii) as defined in Claim 1 wherein x is from about 2.2 to about 4, and wherein R is C_1 - C_8 alkyl and R^1 is methyl, ethyl, \underline{n} -propyl, butyl, \underline{i} -propyl, \underline{n} -octyl, 2-ethylhexyl, \underline{n} -octadecyl, or a mixture of any of the foregoing.
- 4. An oligomer of a triaminotriazine compound (iii) (i) as defined in Claim 1 wherein R is C_1 - C_8 lower alkyl and R^1 is methyl, ethyl, <u>n</u> propyl, butyl, <u>i</u>-propyl, <u>n</u> octyl, 2-ethylhexyl, <u>n</u>-octadecyl, or a mixture of any of the foregoing.
 - 5. A curable composition comprising:
 - (a) an active hydrogen-containing material;
 - (b) a triazine compound selected from
 - (i) a triaminotriazine compound of the formula $C_3N_6(CH_2OR)_{6-x}$ (CH2NHCOOR¹)_x;
 - (ii) a benzoguanamine compound of the formula $C_3N_5(C_6H_5)(CH_2OR)_{4-y}(CH_2NHCOOR^1)_y$;

- (iii) an oligomer of (i) or of (ii); and
- (iv) a mixture of at least two of any of (i), (ii) and (iii), wherein the R groups are, independently, hydrogen or alkyl from 1 to 12 carbon atoms, the R¹ groups are, independently, alkyl of from 1 to 20 carbon atoms, x is in the range of from about 2 to about 6, and y is in the range of from about 2 to about 4; and
 - (c) a cure catalyst.
- 6. A curable composition as defined in Claim 5 wherein the active hydrogen-containing material is a polymeric material containing at least one class of reactive groups selected from carboxyl groups, alcoholic hydroxy groups, amide groups, amine groups or a mixture of any of such groups or a group convertible to any of such groups.
- 7. A curable composition as defined in Claim 5 wherein the triazine compound is a triaminotriazine compound (i) wherein x is from about 2.8 to about 6, R is C_1 - C_8 alkyl and R^1 is methyl, ethyl, n-propyl, butyl, i-propyl, n-octyl, 2-ethylhexyl, n-octadecyl, or a mixture of any of the foregoing.
- 8. A curable composition as defined in Claim 5 wherein the cure catalyst comprises tetrabutyldiacetoxy stannoxane, dibutyltin dilaurate or dimethyltin dilaurate.
- 9. A substrate protectively coated with .a cured composition as defined in Claim 5.
- 10. An article of manufacture comprising a cured composition as defined in Claim 5 and a filler.

